



RNAspace.org: An integrated environment for the prediction, annotation, and analysis of ncRNA.

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RNAspace.org: An integrated environment for the prediction, annotation, and analysis of ncRNA

MARIE-JOSÉE CROS,¹ ANTOINE DE MONTE,² JÉRÔME MARIETTE,³ PHILIPPE BARDOU,⁴
BENJAMIN GRENIER-BOLEY,² DANIEL GAUTHERET,⁵ HÉLÈNE TOUZET,^{2,6}
and CHRISTINE GASPIN^{1,3,6}

¹INRA, UBIA, UR 875, F-31320 Castanet-Tolosan, France

²LIFL, UMR CNRS 8022 Université Lille 1 and INRIA Lille Nord Europe, 59655 Villeneuve d'Ascq cedex, France

³INRA, Plateforme Bioinformatique, F-31320, UR 875, Castanet-Tolosan, France

⁴INRA, SIGENAE, UMR 444, F-31320 Castanet, France

⁵IGM UMR 8621 CNRS-U Paris Sud, 91405 Orsay cedex, France

ABSTRACT

The annotation of noncoding RNA genes remains a major bottleneck in genome sequencing projects. Most genome sequences released today still come with sets of tRNAs and rRNAs as the only annotated RNA elements, ignoring hundreds of other RNA families. We have developed a web environment that is dedicated to noncoding RNA (ncRNA) prediction, annotation, and analysis and allows users to run a variety of tools in an integrated and flexible manner. This environment offers complementary ncRNA gene finders and a set of tools for the comparison, visualization, editing, and export of ncRNA candidates. Predictions can be filtered according to a large set of characteristics. Based on this environment, we created a public website located at <http://RNAspace.org>. It accepts genomic sequences up to 5 Mb, which permits for an online annotation of a complete bacterial genome or a small eukaryotic chromosome. The project is hosted as a Source Forge project (<http://rnaspace.sourceforge.net/>).

Keywords: bioinformatics; noncoding RNA; prediction; annotation

INTRODUCTION

Noncoding RNAs (ncRNA) are RNAs that are transcribed, but not translated into protein. They include well-characterized transfer RNAs and ribosomal RNAs, snRNAs, snoRNAs, and miRNAs, as well as a plethora of new ncRNAs that have been shown to play major roles in the cellular processes of all living organisms (Amaral et al. 2008; Jacquier 2009; Ponting et al. 2009; Waters and Storz 2009; Liu and Carnilli 2010). Even though a large number of genomes have now been sequenced, the number and the diversity of ncRNAs remain largely unknown. The existence of pervasive conserved elements and the extensive expression of transcripts in the noncoding regions of genomes suggest that an important number of ncRNAs of unknown function and structure remain to be identified (Hüttenhofer et al. 2005). In Eukaryotes, hundreds of thousands of (often short) noncoding

transcripts are expressed from the intergenic regions, introns and antisense strands of protein-coding genes (Carninci et al. 2005; van Bakel et al. 2010). Pervasive transcription also occurs in Bacteria (Toledo-Arana et al. 2009) and Archaea (Jäger et al. 2009). Thousands of complete genomes are currently being sequenced, and many more are forthcoming. In such a context of data accumulation, the ability to differentiate automatically known RNA families and the development of additional computational tools dedicated to the improvement of the completeness of the catalog of functional elements are required.

In spite of considerable improvements in the development of ncRNA detection software in the recent years, the prediction and annotation of ncRNAs still remain challenging tasks (Menzel et al. 2009). The most effective ncRNA detection software are distributed as stand-alone applications (Rivas and Eddy 2001; Klein and Eddy 2003; Livny et al. 2005; Washietl et al. 2005), some of them address only specific classes of RNAs (Lowe and Eddy 1997; Laslett and Canbäck 2004; Schattner et al. 2006; Lagesen et al. 2007) or organisms (Klein et al. 2002; Schattner 2002; Upadhyay et al. 2005; Larsson et al. 2008). The integration of these approaches for consistent ncRNA prediction and annotation

⁶Corresponding authors.

E-mail christine.gaspin@toulouse.inra.fr.

E-mail helene.touzet@lifl.fr.

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remains the realm of bioinformatics experts. As a result, ncRNA annotation in genome databases lags behind protein-coding gene annotation. This is also due in large part to the relative difficulty of identifying ncRNA-encoding loci, which requires unique bioinformatic approaches. Integrating the diversity of ncRNA prediction and annotation tools in a unique environment is highly desirable for biologists interested in the prediction, annotation, and basic analysis of known and new ncRNA genes from genomic sequences.

The most ambitious attempt at providing tools for RNA prediction and annotation is that of the RFAM database (Gardner *et al.* 2009). RFAM 10.0 hosts over 1446 ncRNA families and allows biologists to perform similarity searches against this database. Users can also access precomputed full-genome annotations. The limitations of the RFAM website are the small size limit for online searches and the restriction of matches to existing RNA families, which excludes a large number of ncRNAs for which no alignment is available or submitted to RFAM. Other ncRNA databases exist that focus on a single class of molecules such as miRBase (Griffiths-Jones *et al.* 2008), on one class for a set of organisms such as the snoRNABase (Lestrade and Weber 2006), or on one genome such as ASRP (<http://asrp.cgrb.oregonstate.edu/db/>) or that aim at providing an exhaustive catalog of ncRNAs such as NONCODE (Liu *et al.* 2005) or fRNAdb (Kin *et al.* 2006). Besides RFAM, several alternative computational methods exist that perform ncRNA prediction in genomic sequences. These can be classified into two types, namely, those that aim at searching for ncRNA homologs using conserved sequence and structure characteristics and those that aim at discovering new ncRNA families. Methods of the former type use sequence/structure alignments in order to identify key conserved motifs involved in the molecule structure. Alignments are then processed and modeled in several ways, for instance, with position weight matrices in ERPIN (Gautheret and Lambert 2001) or covariance models in INFERNAL (Nawrocki *et al.* 2009). These types of methods also include programs dedicated to one type of ncRNA, such as tRNAScan-SE (Lowe and Eddy 1997) and RNAMmer (Lagesen *et al.* 2007). Programs such as RNAMOTIF (Macke *et al.* 2001), PatScan (Dsouza *et al.* 1997), and DARN! (Zytnicki *et al.* 2008) provide users with a programming language allowing the description of any ncRNA structure. Methods of the latter type, namely, programs aiming at *de novo* ncRNA identification, use diverse and often complementary approaches, including sequence composition bias analysis and the identification of significant stable and conserved secondary structure regions in multiple sequence alignments. Successful ncRNA searches using these approaches have been reported in Archaea (Klein *et al.* 2002; Schattner 2002), Bacteria (Rivas *et al.* 2001; Livny *et al.* 2006; Geissmann *et al.* 2009), and Eukaryotes (Mendes *et al.* 2009; Li *et al.* 2010). More recently, the CMfinder tool (Yao *et al.* 2006) was used to predict large sets of new ncRNA/motifs signatures in Bacteria (Weinberg *et al.* 2007, 2009).

The absence of a genuine generic ncRNA prediction method and the expertise required to detect ncRNAs based on sequence and structure conservation have led us to develop a web environment, RNAspace.org, that permits users to perform all of these tasks in an integrated fashion. This web environment is dedicated to biologists who are involved in genome annotation projects with a particular interest in known and new ncRNA families. RNAspace.org enables these users to run a variety of ncRNA gene finders in an integrated way and to explore results using dedicated tools for the comparison, visualization, filtering, alignment, and editing of putative ncRNAs. This environment is collaboratively developed and uses open-source software. Thus, computer scientists developing new methods on these topics may be interested in using the web environment to integrate their methods.

RESULTS

The ncRNA prediction and annotation activities are organized into three main steps corresponding to a typical annotation process. In the first step, users upload the sequence(s) to be analyzed, along with related information. In the second step, users select ncRNA gene finders. It will be possible to return to this step later on for selecting other gene finders or changing their parameters. The third step includes a set of functionalities to assess predicted ncRNA genes with regards to their context, annotation, conservation, and secondary structure. Results obtained from an analysis can be saved in various standard formats and are kept 14 d.

Data upload step

The sequences to be analyzed are entered as a fasta or multi-fasta file. One or more sequence files can be uploaded, provided that the total length is <5 Mb. As some software require knowledge of the phylogenetic domain (Archaea, Bacteria, Eukarya) of the organism under analysis, this information must be entered. Other optional information can be entered. An e-mail address is required in order to receive the link giving access to the results of the prediction step.

ncRNA prediction step

After sequence uploading, users select ncRNA gene finders. These gene finders were chosen so that a variety of complementary approaches can be used, modeling different characteristics of ncRNAs. Programs are organized into three main groups: homology search, comparative analysis, and *ab initio* prediction. The first class of tools identifies genes that belong to known families of ncRNAs, whereas the two latter classes aim at finding new families. The list of tools currently available is presented in Table 1.

TABLE 1. List of software available in the predict step

Software (reference)	Used for
Homology search tools	
BLAST (Altschul et al. 1990)	Sequence similarity search against ncRNA databases
YASS (Noé and Kucherov 2005)	Sequence similarity search against ncRNA databases
Infernal (Nawrocki et al. 2009)	RNA signature search (primary+secondary structure)
Erpin (Gautheret and Lambert 2001)	RNA signature search (primary+secondary structure)
Darn! (Zytnicki et al. 2008)	RNA signature search (primary+secondary structure)
RNAmmer (Lagesen et al. 2007)	Search for ribosomal RNAs
tRNAscan-SE (Lowe and Eddy 1997)	Search for transfer RNAs
Comparative analysis	
Step 1: pairwise sequence alignment	
BLAST (Altschul et al. 1990)	Sequence similarity
YASS (Noé and Kucherov 2005)	Sequence similarity
Step 2: identification of conserved regions across species	
CG-seq	Combination of pairwise alignments into clusters of conserved sequences
Step 3: looking for a conserved consensus secondary structure	
CaRNAC (Touzet and Perriquet 2004)	Secondary structure prediction
RNAz/clustalW (Washietl et al. 2005; Larkin et al. 2007)	Secondary structure prediction
Ab initio prediction	
AtypicalGC	Nucleotide bias composition

Homology search tools rely on the knowledge of a signature for a given ncRNA family. This signature contains information at the sequence and/or secondary structure level. For sequence level searches, RNAspace compares user sequences to several databases using the sequence alignment programs BLASTN (Altschul et al. 1990) and YASS (Noé and Kucherov 2005). Both programs implement a heuristic approach for local similarity search. BLASTN is based on contiguous k-mers, while YASS is based on spaced-seeds that are known to achieve higher sensitivity. However, BLASTN is faster than YASS. When aligning the query sequence against all sequences of a given ncRNA database with BLASTN or YASS, it appears frequently that the query sequence matches against many sequences from the same family in the database, giving rise to a high number of alignments. Of course, all these alignments correspond to a single RNA gene on the query sequence. So all alignments referring to the same region on the same strand for a same family are merged in a single prediction provided that the positions between the alignments are consistent. We say that two local alignments A and A' are consistent if $|(a - x) - (a' - y)| < 10$, where a is the start position of the query sequence in A , a' is the start position of the query sequence in A' , x is the start position of the ncRNA sequence of the family in A , and y is the start

position of the ncRNA sequence of the family in A' . Alignments are merged in a greedy fashion, starting from the first alignment on the query sequence, from 5' to 3'. Among all mutually consistent alignments, only the best alignments (with lower P -value) are stored and can be presented in a visual format to the user. The number of stored alignments is parameterizable. The start (respectively, end) position of the predicted ncRNA is obtained as the lower (respectively, upper) bound of start (respectively, end) positions of alignments on the query sequence. Installed search databases are currently RFAM 9.1 (1371 families) and RFAM 10.0 (1446 families less the SSU_rRNA_5 family) (Gardner et al. 2009), fRNAdb 3.0 (Kin et al. 2006), and miRBase 13.0 (Griffiths-Jones et al. 2008). For secondary structure and sequence-based homology searches, users may run ERPIN (Gautheret and Lambert 2001), INFERNAL (Nawrocki et al. 2009), and DARN! (Zytnicki et al. 2008). These programs use RNA secondary structure profiles. INFERNAL uses covariation models, which are very sensitive profiles, and offers all RFAM 10.0 families. It is possible to launch it

on all families simultaneously or to select one family at a time. Almost all RFAM 9.1 families are currently modeled in ERPIN, and the underlying search algorithm is faster than INFERNAL. DARN! provides specific patterns for C/D box sRNA, some riboswitches, and tRNA. Note that these programs are significantly slower than similarity sequence search programs but are able to identify homologous ncRNA with a lower sequence similarity to known ncRNA. Finally, RNAspace users may run specific programs for rRNA and tRNA gene prediction, respectively, RNAmmer (Lagesen et al. 2007) and tRNAscan-SE (Lowe and Eddy 1997), which are both recognized for their high prediction accuracy.

RNAspace users may also identify new ncRNAs with no homology with the known ncRNAs, using comparative sequence analysis. The underlying principle in this approach is that functional ncRNAs are under a positive selection pressure, hence their sequence should be better conserved than random noncoding regions. Furthermore, mutations observed between homologous structured RNA sequences should be consistent with the formation of a conserved secondary structure core. RNAspace includes the basic components required for running a comparative analysis pipeline. For bacterial genomes, the predictions obtained with our comparative pipeline will differ somewhat from those made by the Livny group (Livny et al. 2008), who

used the presence of a transcription terminator, and not secondary structure conservation, as the main filtering criteria. Our comparative analysis results should resemble more those obtained in recent studies combining conservation analysis and the RNaz filter, such as that by Sonnleitner et al. (2008). To run the comparative prediction tool, users select a set of species for comparisons (at most four species). Due to the relatively long computer runtimes involved, we limited the comparative analysis to Bacteria and Archaea. In the first component of the pipeline, the query sequence is compared against all intergenic regions of the selected organisms by selecting BLASTN or YASS. The expected result is a set of similar sequences across species. The second component, CG-seq (<http://bioinfo.lifl.fr/CGseq/>), performs the combination of all pairwise alignments into clusters of significantly conserved regions. Finally users may select caRNAC (Touzet and Perriquet 2004) or RNaz (Washietl et al. 2005) as the third pipeline component, in charge of inferring and scoring the conserved consensus secondary structure. RNaz requires pre-aligned sequences (a ClustalW [Larkin et al. 2007] alignment is first performed by RNAspace) and is known to be well suited to process clusters of highly similar sequences. caRNAC is a simultaneous align/fold algorithm that shows a better specificity when sequences are difficult to align accurately (Gardner and Giegerich 2004).

The third group of tools performs *ab initio* prediction by seeking intrinsic signals, which aid in distinguishing ncRNA from other elements in the genome. In the past, several attempts were made to characterize ncRNA using folding measures alone (Freyhult et al. 2005). However, none provided a reliable ncRNA prediction at the genome scale. We chose instead to implement a tool that exploits the compositional bias between ncRNA and the rest of the genome. This approach succeeded at predicting ncRNA from (GC)-atypical regions in hyperthermophile archaea (Klein et al. 2002; Schattner 2002). We implemented a method for identifying (GC)-atypical regions in the program AtypicalGC. This program searches for rich (default parameter) atypical GC regions by using a sliding window. Only regions having a GC value distant of 2 SDs from the mean computed on the whole-genomic sequence are considered as atypical.

Users may select a selection of prediction programs simultaneously to obtain a ncRNA catalog that is as complete as possible, given the query sequence. All programs can be run with default parameter values. For most of them, however, parameters can be easily varied through a dedicated interface, according to known biological constraints or user expertise. The analysis performed with a given tool selection is called a *run*. The set of putative ncRNA genes obtained in a *run* may contain redundant overlapping predictions, thus representing accumulated evidence from different ncRNA gene finders. These redundant predictions can be partially merged, given they are consistent in location, strand, length, and functional annotation. Redundancy exists when at least two ncRNA gene finders produce

candidates that overlap on the same strand and share the same functional annotation. Overlapping is considered if putative RNAs share at least 10 nucleotides. Functional annotation is based on the family name. For that purpose, a table of synonymous names has been built by hand. It gives an equivalence between name families assigned by different software. Predictions share the same functional annotation if they possess identical or synonymous names according to this equivalence table. Redundancy is managed using an algorithm that merges regions by considering positions, functional annotation, and the nature of the priority assigned to gene finders. Because accuracy of some family-dedicated gene finders is now well established, redundancy can be removed by giving them higher priority for assigning begin/end positions. In RNAspace, tRNAscan-SE and RNAmmer were selected as such tools. Finally, for merging ncRNAs, the algorithm first considers the priority assigned to selected gene finders and then the positions to be merged. This rule does not hold for ncRNAs, which are added by the user, and for ncRNAs annotated as *unknown* (such as those provided by the comparative analysis pipeline or with *ab initio* approach). In practice, this merging strategy provides a valuable way to deal with redundant predictions and simplifies the analysis of predictions. Note, however, that in some cases the algorithm is not always a satisfying solution. The latter case is illustrated by a simple example when two tRNAs distant of a few nucleotides are predicted by tRNAscan-SE as two separated regions and are also predicted by BLASTN or YASS as a single one. In this case, no merging is realized by the current merging functionality.

Explore step

Different gene finders will typically differ in the precise boundaries and annotations of a putative ncRNA. In this case, RNAspace aims to display as much information as possible to help users decide on a suitable selection of evidence and prediction. This explore step provides users with a set of information, functionalities, and programs dedicated to the detailed analysis of predictions and to data export. Prediction results can be visualized as a summary table showing the main characteristics of each hit, or using one of the Jbrowse (Skinner et al. 2009) or CGview (Stothard and Wishart 2005) graphical visualization tools (Fig. 1). Under the table view, users can dynamically sort and explore ncRNA predictions. Users may also apply filtering criteria to any data field by comparing its value to a regular expression. Only predictions matching all user-defined filters are displayed. By default, results are ranked according to their physical location, but any other data field (name, software, etc.) can be used as a sort criterion. Each line of the table shows a ncRNA prediction and offers access to a set of functionalities and data, such as the sequence context of the prediction, existing secondary

structures, and alignments in which the sequence is involved (Fig. 2). Other information available for each prediction includes the name of the user sequence, the RNA family (when applicable), the start and end positions on the user sequence, the strand on which the prediction was made (when possible), the domain of life, species and a replicon of the user sequence, the program or set of programs that produced the prediction, the number of

alignments in which the prediction is involved, and the identification of the run in which the prediction was performed. Alignments come from two sources. They can be the result of prediction by sequence similarity search. In this case, they are the best pairwise alignments computed by BLASTN and YASS. Otherwise, they consist of multiple alignments computed online from a (multiple) selection of predictions.

A

RNAspace.org
The non-coding RNA annotation platform

Home 1. Load data 2. Predict 3. Explore HELP

Current results for the 001940b5dc7db2b project: **31 putatives RNAs predicted.**

Software tools used and user actions are summarized in the left re-sizeable table and query sequence(s) in the right re-sizeable table. See the project [history](#) for more details.

Run or user identifier	Description	Number of RNAs
r01	BLAST/Rfam_10.0_seed	19
	RNAmer	6
	tRNAscan-SE	6

Query sequence(s)
sample 100001 nt bacteria E.coli K12 Chromosome

Field Operator Value (wildcards allowed) Result
Criterium Comparison Give value Add/Update 31/31 RNAs satisfy filter(s)

Table view JBrowse view CGview view

The table of results may be sorted by clicking on the column titles. You can select predictions by ticking the check boxes in the left column and perform actions on them using the down-drop lists below the table [7].

Predictions 1 - 20 of 31 Display Terse set Show 20 Page 1 of 2 >>

All	ID	Seq name	Family	Start	End	Size	Strand	Species	Domain	Replicon	Software	Align	Run
<input type="checkbox"/>	000014	sample	Cobalamin	4991	5181	191	+	E.coli	bacteria	Chromosome	BLAST/Rfam_10.0_seed	5	r01
<input type="checkbox"/>	000011	sample	16s_rRNA	8273	9802	1530	+	E.coli	bacteria	Chromosome	RNAmer	0	r01
<input type="checkbox"/>	000001	sample	tRNA-Glu	9979	10054	76	+	E.coli	bacteria	Chromosome	tRNAscan-SE	0	r01
<input type="checkbox"/>	000016	sample	tRNA	9979	10051	73	+	E.coli	bacteria	Chromosome	BLAST/Rfam_10.0_seed	5	r01
<input type="checkbox"/>	000007	sample	23s_rRNA	10250	13151	2902	+	E.coli	bacteria	Chromosome	RNAmer	0	r01
<input type="checkbox"/>	000022	sample	PK-G12rRNA	12530	12637	108	+	E.coli	bacteria	Chromosome	BLAST/Rfam_10.0_seed	5	r01
<input type="checkbox"/>	000030	sample	5S_rRNA	13246	13361	116	+	E.coli	bacteria	Chromosome	BLAST/Rfam_10.0_seed	5	r01
<input type="checkbox"/>	000009	sample	5s_rRNA	13248	13362	115	+	E.coli	bacteria	Chromosome	RNAmer	0	r01
<input type="checkbox"/>	000024	sample	Trp_leader	13332	13385	54	+	E.coli	bacteria	Chromosome	BLAST/Rfam_10.0_seed	1	r01
<input type="checkbox"/>	000002	sample	tRNA-Thr	16995	17070	76	+	E.coli	bacteria	Chromosome	tRNAscan-SE	0	r01
<input type="checkbox"/>	000017	sample	tRNA	16995	17067	73	+	E.coli	bacteria	Chromosome	BLAST/Rfam_10.0_seed	5	r01
<input type="checkbox"/>	000003	sample	tRNA-Tyr	17079	17163	85	+	E.coli	bacteria	Chromosome	tRNAscan-SE	0	r01
<input type="checkbox"/>	000018	sample	tRNA	17079	17160	82	+	E.coli	bacteria	Chromosome	BLAST/Rfam_10.0_seed	2	r01
<input type="checkbox"/>	000026	sample	Rit	17090	17148	59	+	E.coli	bacteria	Chromosome	BLAST/Rfam_10.0_seed	2	r01
<input type="checkbox"/>	000004	sample	tRNA-Gly	17280	17354	75	+	E.coli	bacteria	Chromosome	tRNAscan-SE	0	r01
<input type="checkbox"/>	000019	sample	tRNA	17280	17351	72	+	E.coli	bacteria	Chromosome	BLAST/Rfam_10.0_seed	5	r01
<input type="checkbox"/>	000005	sample	tRNA-Thr	17361	17436	76	+	E.coli	bacteria	Chromosome	tRNAscan-SE	0	r01
<input type="checkbox"/>	000020	sample	tRNA	17361	17433	73	+	E.coli	bacteria	Chromosome	BLAST/Rfam_10.0_seed	5	r01
<input type="checkbox"/>	000028	sample	TPP	21269	21352	84	+	E.coli	bacteria	Chromosome	BLAST/Rfam_10.0_seed	1	r01
<input type="checkbox"/>	000027	sample	P26	22537	22598	62	+	E.coli	bacteria	Chromosome	BLAST/Rfam_10.0_seed	5	r01

With selected predictions: Edit... Analyse... Export... With all predictions: EXPORT...

Comments and remarks: contact@rnaspace.org

FIGURE 1. (Continued on next page)

Users may edit most of the information associated to a prediction, produce and save alternative secondary structures computed with one of the RNAfold (Gruber *et al.* 2008), UNAFold (Markham and Zuker 2008), or RNASHapes (Steffen *et al.* 2006) programs, or provide their own secondary structure in the form of balanced parentheses. The structures

are generated with RNAplot (Gruber *et al.* 2008) or VARNA (Darty *et al.* 2009). Users may also select one or more predictions in the table and apply editing functionalities to them. Thus, users can modify the RNA family name, create a new prediction from the selection, delete the predictions, or split the predictions into two new families. Users may also

B

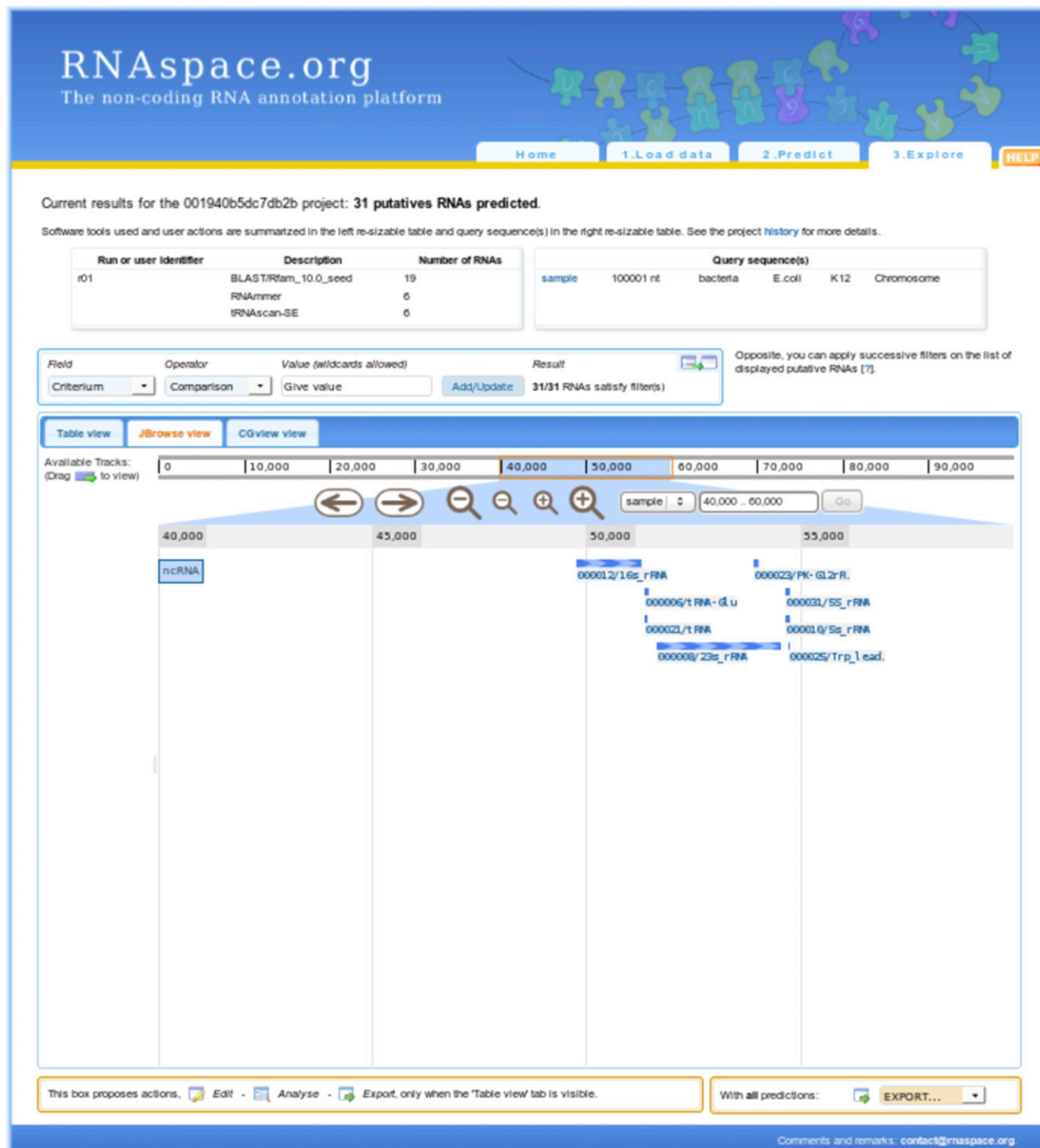


FIGURE 1. (Continued on next page)

analyze selected predictions. A functionality allows for the alignment of a selection of predictions on the basis of their sequence and structure conservation. The alignment is built with ClustalW (Larkin et al. 2007), and the common structure is derived with RNAz (Washietl et al. 2005). It is also possible

to map and visualize a selection of predictions in ApolloRNA (<http://carlit.toulouse.inra.fr/ApolloRNA>), an extension of the annotation environment Apollo (Lewis et al. 2002) dedicated to RNA analysis. Last, selected predictions can be exported under the formats multifasta, gff, RNAML, and CSV.

C

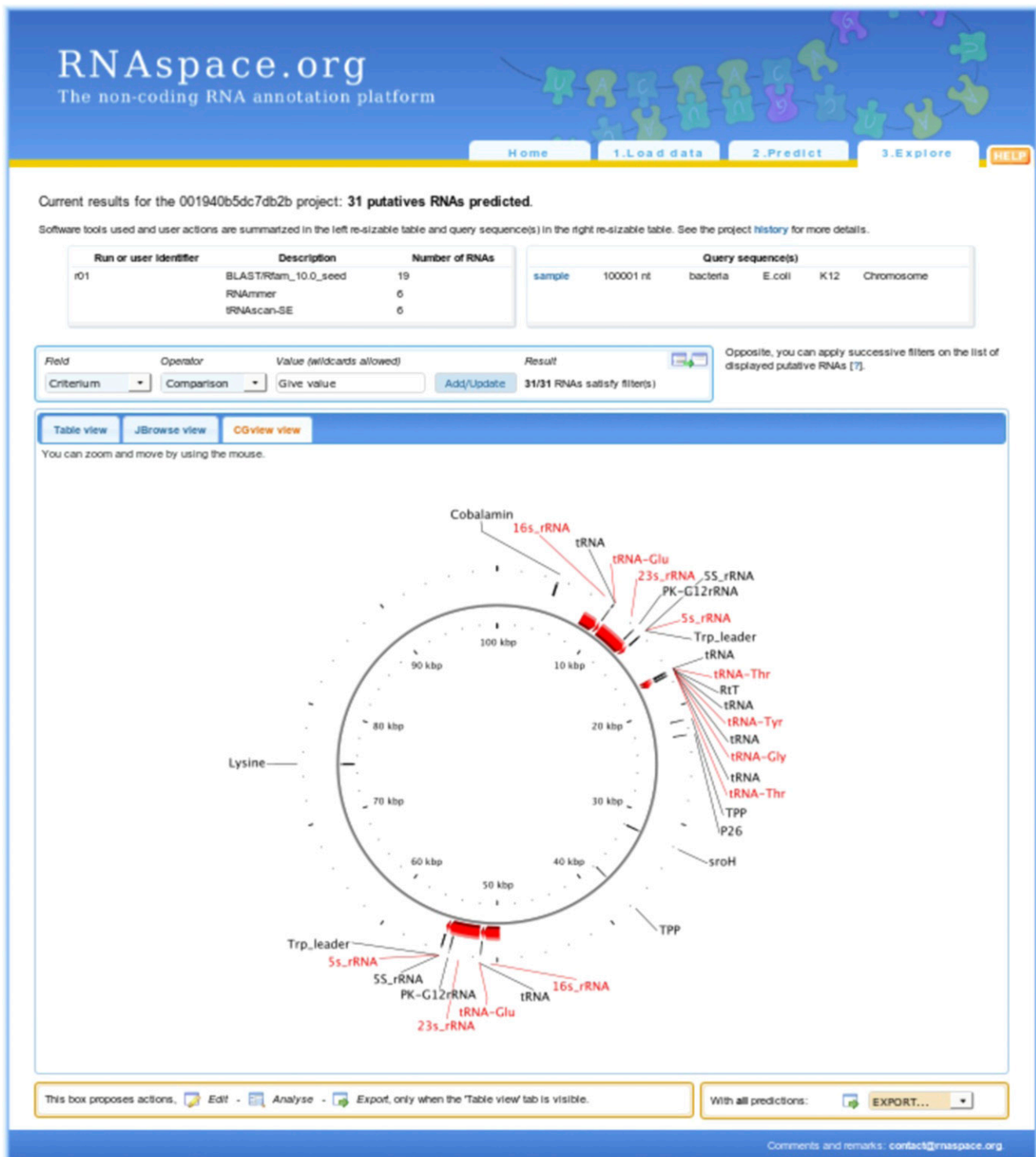


FIGURE 1. Visualization of predictions in the table view (A), with JBrowse (B), and with CGview (C).

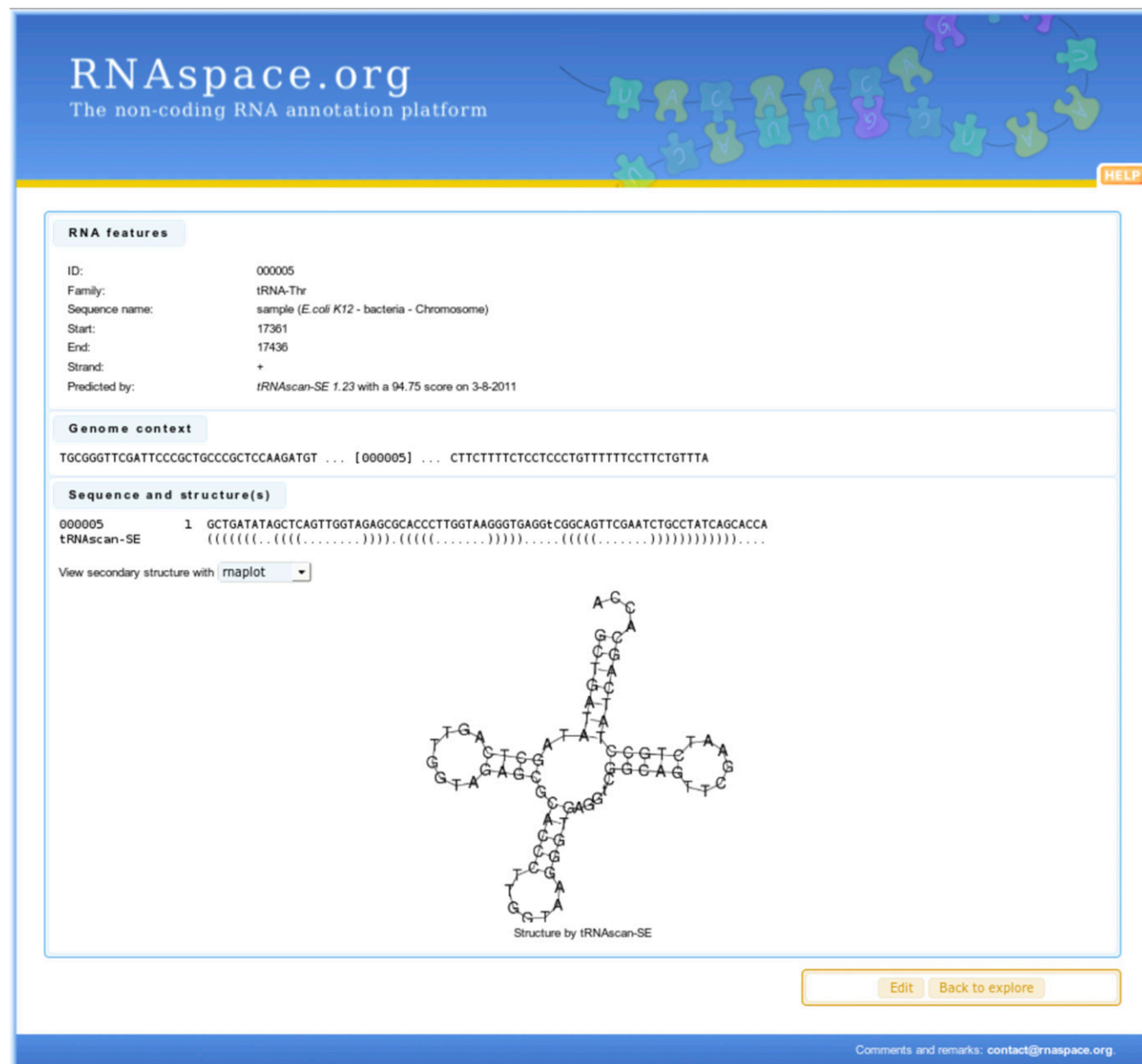


FIGURE 2. Visualization of the characteristics of a predicted ncRNA.

DISCUSSION

The annotation of ncRNA remains a major bottleneck on the way toward understanding genome function and evolution. In this context, RNAspace.org provides to biologists a user-friendly web environment dedicated to ncRNA prediction, annotation, and analysis, allowing users to run a variety of complementary tools in an easy and integrated way. Biologists involved in the annotation of ncRNAs in any newly sequenced genome can predict ncRNAs from known families and explore potential novel ncRNAs using different tools. In archaeal or bacterial genomes, users may

also discover new ncRNA families using comparative genomics. RNAspace.org does not yet offer access to all the programs and functionalities that may be desirable for ncRNA annotation. We first integrated those tools and databases that partner laboratories had contributed to develop or were used to working with.

In the future, we plan to develop the environment along several lines. First, it will be necessary to extend the scope of ncRNA predictions by including, for example, dedicated CRISPR and miRNA gene finders. Second, as observed in our tests, the combination of gene finders may produce large prediction sets, making it difficult for users to focus

on the most significant predictions. A functionality is already implemented to help users reduce redundancy by merging overlapping regions. This functionality will be improved in the future. For instance, the automatic merging of overlapping predictions should benefit strongly from a continuous updating of the dictionary of synonymous RNA gene names. Other functionalities are implemented to help users edit structural and functional annotations and merge or split annotated RNA families. These functionalities will be further developed and improved. For example, we plan to allow for the visualization of predictions in the context of other genome annotation (such as protein-coding gene data). This information will improve ncRNA prediction through knowledge of neighboring gene functions and of the intronic, coding, or intergenic locations of candidate ncRNA. We will also consider the integration of existing tools for editing multiple alignments to enable expert users to improve the accuracy of multiple alignments for consensus structure identification.

With the huge quantity of high-throughput sequencing data obtained by transcriptome studies, it is also highly desirable to consider RNAseq and sRNAseq data. In the future, we will consider handling large sets of RNAseq and sRNAseq with respect to a reference sequence. Such analyses will permit both the annotation and quantification of noncoding transcripts and the search for potential targets of regulatory RNA acting through RNA–RNA interactions.

RNAspace is a collaborative project, and our goal is to make this environment available to the largest community. By making the code open-source, we hope that useful features of RNAspace will be adapted and extended by other developers to serve the needs of particular user groups.

MATERIALS AND METHODS

Implementation and architecture

RNAspace is developed in Python. The web framework CherryPy and the template engine Cheetah were chosen for their simplicity and proximity with Python. A three-tiers client-server architecture is implemented with a presentation tier that displays information, an application tier that controls application's functionality by performing processing, and a data tier (data access) that keeps data independent from the application tier. The Presentation tier relies on a Model-View-Controller (MVC) design pattern. The model is related to domain-specific representation of the information on which the application operates (here, process and access data). The view renders the model into a form suitable for interaction, typically a user interface element (here, template). The controller processes and responds to events, typically user actions, and may invoke changes on the model (here, view as web request entry). The source code is developed under the GPL open-source license. The project is hosted as a Source Forge project (<http://sourceforge.net/projects/rnaspace>). More information on the architecture and development is available in the RNAspace development documentation.

Data storage

RNAML (Waugh et al. 2002) is used to store and export information on putative ncRNA. RNAML is a standard XML syntax for RNA information. This format is used as an input or output format in a variety of applications (Ruan et al. 2004; Jossinet and Westhof 2005; Noirot et al. 2008; Darty et al. 2009). We choose RNAML to store data produced all along an analysis because it provides a single format for representing information specific to RNA molecules and it proposes a standard syntax to easily express information on ncRNA, thus offering a way to represent a large amount of information in an unambiguous and reusable syntax.

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